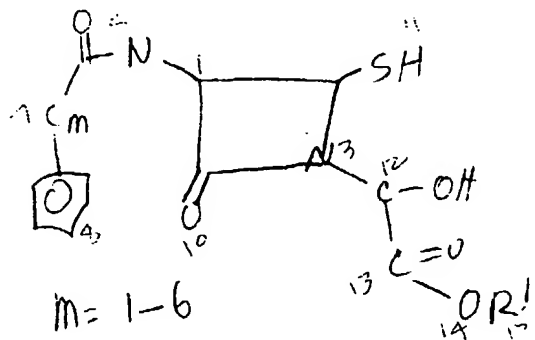


SEARCH REQUEST FORM

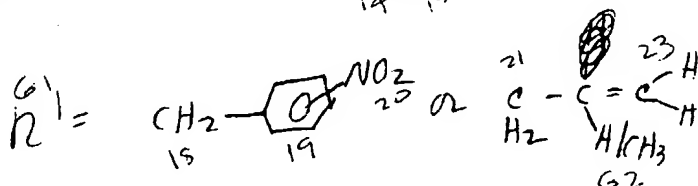
Requestor's Name: Bent Serial Number: 10/1006579
 Date: 10/3 Phone: 4105 978 Art Unit: 162
10/1781188

Search Topic:

Please write a detailed statement of search topic. Describe specifically as possible the subject matter to be searched. Define any terms that may have a special meaning. Give examples or relevant citations, authors, keywords, etc., if known. For sequences, please attach a copy of the sequence. You may include a copy of the broadest and/or most relevant claim(s).



Point of Contact:
Beverly Shears
Technical Info. Specialist
CM1 1E05 Tel: 308-4994



10/1781188

Do Not broaden search

(135)

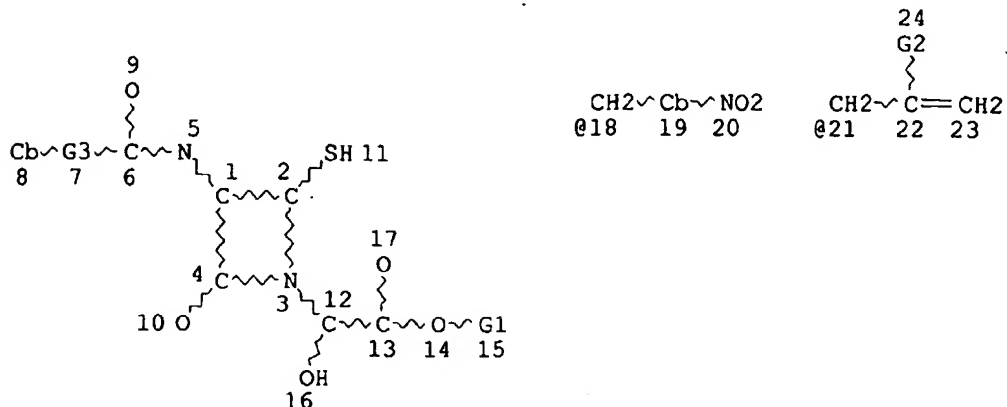
STAFF USE ONLY

Date completed: <u>10-04-02</u>	Search Site	Vendors
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Elapsed time:	<input type="checkbox"/> Pre-S	<input type="checkbox"/> Dialog
CPU time:	<input type="checkbox"/> Type of Search	<input type="checkbox"/> APS
Total time: <u>24</u>	<input type="checkbox"/> N.A. Sequence	<input type="checkbox"/> Geninfo
Number of Searches:	<input type="checkbox"/> A.A. Sequence	<input type="checkbox"/> SDC
Number of Databases: <u>1</u>	<input type="checkbox"/> Structure	<input type="checkbox"/> DARC/Questel
	<input type="checkbox"/> Bibliographic	<input type="checkbox"/> Other

Berch
10/006579

10/006579

L7 (FILE 'REGISTRY' ENTERED AT 15:38:11 ON 04 OCT 2002)
STR



VAR G1=18/21
VAR G2=H/CH3
REP G3=(1-6) C
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
GGCAT IS UNS AT 8
DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:
RSPEC I
NUMBER OF NODES IS 24

STEREO ATTRIBUTES: NONE
569- NSEA FILE=REGISTRY SSS FUL L7

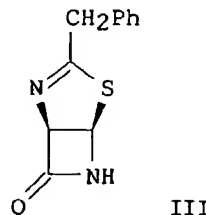
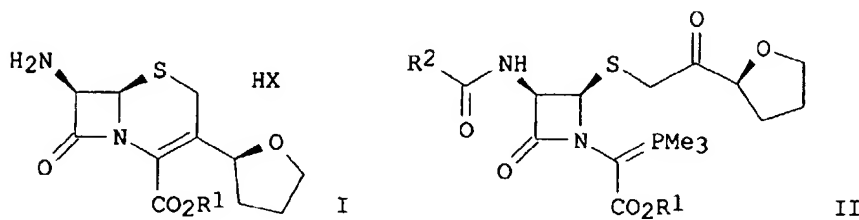
100.0% PROCESSED 165 ITERATIONS 1 ANSWERS
SEARCH TIME: 00.00.03

L10 FILE 'HCAPLUS' ENTERED AT 15:45:34 ON 04 OCT 2002
1 S L9

L10 ANSWER 1 OF 1 HCAPLUS COPYRIGHT 2002 ACS
ACCESSION NUMBER: 2002:449689 HCAPLUS
DOCUMENT NUMBER: 137:33162
TITLE: Process for the preparation of p-nitrobenzyl or
allyl esters of 3-cyclic-ether substituted
cephalosporins from trimethylphosphinic
compounds via an intramolecular Wittig reaction
INVENTOR(S): Colberg, Juan Carlos; Tucker, John Lloyd;
Zenoni, Maurizio; Fogliato, Giovanni; Donadelli,
Alessandro
PATENT ASSIGNEE(S): Pfizer Products Inc., USA
SOURCE: PCT Int. Appl., 47 pp.
CODEN: PIXXD2
DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

10/006579

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2002046199	A1	20020613	WO 2001-IB2181	20011119
W:	AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM			
RW:	GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG			
AU 2002023929	A5	20020618	AU 2002-23929	20011119
US 2002099205	A1	20020725	US 2001-6579	20011204
PRIORITY APPLN. INFO.:			US 2000-251018P	P 20001204
			WO 2001-IB2181	W 20011119
OTHER SOURCE(S):			CASREACT 137:33162; MARPAT 137:33162	
GI				



AB A process for the prepn. of I ($R_1 = p$ -nitrobenzyl, allyl; $X = \text{halo}$) via an intramol. Wittig reaction of II ($R_1 = p$ -nitrobenzyl, allyl; $R_2 = C_1\text{-}6\text{-alkyl}, C_6\text{-}10\text{-aryl}, C_6\text{-}10\text{-aryl-}C_1\text{-}6\text{-alkyl}, \text{dithianyl}$) to prep. 3-cyclic-ether substituted derivs. of cephalosporins is described. Thus, III was treated with p -nitrobenzyl glyoxylate monohydrate followed by redn. of the intermediate with NaBH_4 . The resulting hydroxy compd. was treated with p -toluenesulfonic acid followed by addn. of (S)-1-(tetrahydro-2-furanyl)ethanone, addn. of thionyl chloride, and finally trimethylphosphine to give the desired intermediate II ($R_1 = p$ -nitrobenzyl, $R_2 = \text{PhCH}_2$). Cyclization of II via an intramol. Wittig reaction was accomplished by refluxing for 16 h in THF. Addn. of phosphorus pentachloride and α -picoline in dichloromethane gave the free amine of I ($R_1 = p$ -nitrobenzyl).

IT

RL: IMF (Industrial manufacture); RCT (Reactant); SPN (Synthetic

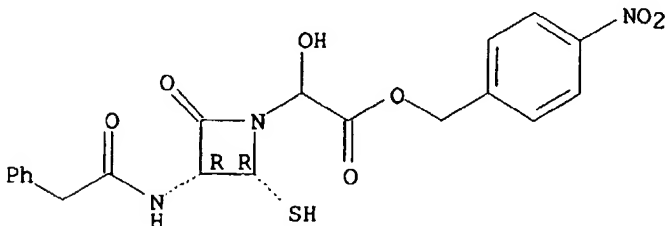
10/006579

preparation); PREP (Preparation); RACT (Reactant or reagent)
(process for the prepn. of p-nitrobenzyl or allyl esters of
3-cyclic-ether substituted cephalosporins from
trimethylphosphinic compds. via an intramol. Wittig reaction)

RN 436800-39-4 HCPLUS

CN 1-Azetidineacetic acid, .alpha.-hydroxy-2-mercapto-4-oxo-3-
[(phenylacetyl)amino]-, (4-nitrophenyl)methyl ester, (2R,3R)- (9CI)
(CA INDEX NAME)

Absolute stereochemistry.



REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR
THIS RECORD. ALL CITATIONS AVAILABLE IN
THE RE FORMAT

L11 FILE 'CAOLD' ENTERED AT 15:46:12 ON 04 OCT 2002
0 S L9

L12 FILE 'USPATFULL' ENTERED AT 15:46:18 ON 04 OCT 2002
1 S L9

L12 ANSWER 1 OF 1 USPATFULL

ACCESSION NUMBER: 2002:186282 USPATFULL
TITLE: Process and ester derivatives useful for
preparation of cephalosporins
INVENTOR(S): Colberg, Juan C., Norwich, CT, UNITED STATES
Tucker, John L., Niantic, CT, UNITED STATES
Zenoni, Maurizio, Milan, ITALY
Fogliato, Giovanni, Bergamo, ITALY
Donadelli, Alessandro, Lodi, ITALY
PATENT ASSIGNEE(S): Pfizer Inc. (U.S. corporation)

	NUMBER	KIND	DATE
PATENT INFORMATION:	US 2002099205	A1	20020725
APPLICATION INFO.:	US 2001-6579	A1	20011204 (10)

	NUMBER	DATE
PRIORITY INFORMATION:	US 2000-251018P	20001204 (60)
DOCUMENT TYPE:	Utility	
FILE SEGMENT:	APPLICATION	
LEGAL REPRESENTATIVE:	PFIZER INC, 150 EAST 42ND STREET, 5TH FLOOR - STOP 49, NEW YORK, NY, 10017-5612	
NUMBER OF CLAIMS:	39	
EXEMPLARY CLAIM:	1	
LINE COUNT:	1433	

Searcher : Shears 308-4994

10/006579

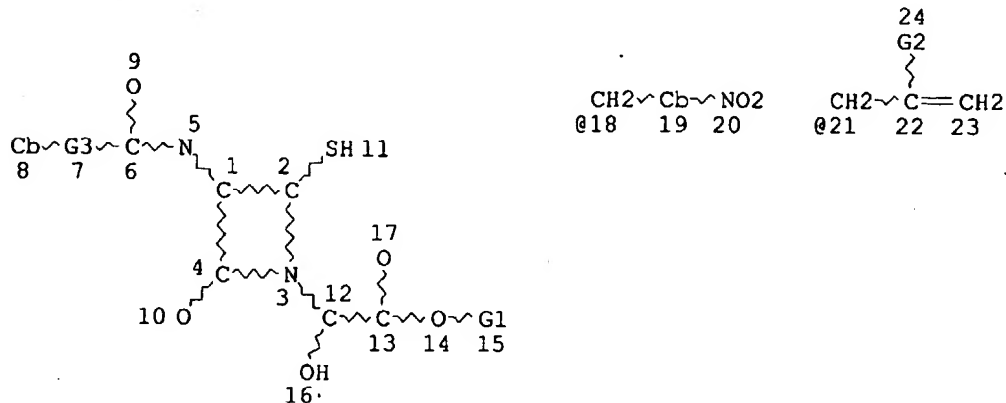
CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB This invention relates a process for preparing a compound of formula (I) ##STR1##

wherein R.sup.1 is para-nitrobenzyl or allyl; and X is halo, which is useful to prepare 3-cyclic-ether-substituted cephalosporins, from trimethylphosphinic compounds. This invention also relates to compounds useful in such process.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L7 (FILE: 'MARPAT' ENTERED AT 15:46:35 ON 04 OCT 2002)
STR



VAR G1=18/21

VAR G2=H/CH3

REP G3=(1-6) C

NODE ATTRIBUTES:

DEFAULT MLEVEL IS ATOM

GGCAT IS UNS AT 8

DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RSPEC I

NUMBER OF NODES IS 24

STEREO ATTRIBUTES: NONE

ATTRIBUTES SPECIFIED AT SEARCH-TIME:

MLEVEL IS CLASS ON RING NODES AND RING GROUPS

MLEVEL IS CLASS ON CHAIN NODES AND CHAIN GROUPS

ECLEVEL IS LIM ON ALL NODES

ALL RING(S) ARE ISOLATED

L14

SEA FILE=MARPAT SSS FUL L7 (MODIFIED ATTRIBUTES)

100.0% PROCESSED 26751 ITERATIONS (6 INCOMPLETE) 7 ANSWERS
SEARCH TIME: 00.02.02

L14 ANSWER 1 OF 7 MARPAT COPYRIGHT 2002 ACS

ACCESSION NUMBER: 137:33162 MARPAT

TITLE: Process for the preparation of p-nitrobenzyl or

Searcher : Shears 308-4994

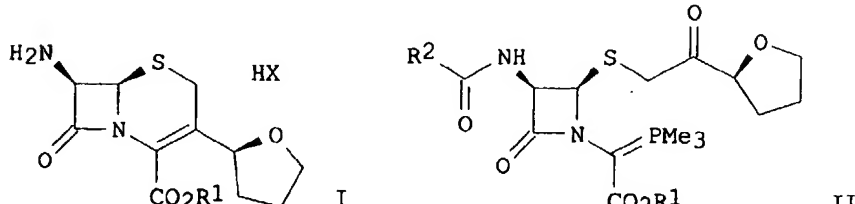
10/006579

allyl esters of 3-cyclic-ether substituted cephalosporins from trimethylphosphinic compounds via an intramolecular Wittig reaction

INVENTOR(S): Colberg, Juan Carlos; Tucker, John Lloyd;
Zenoni, Maurizio; Fogliato, Giovanni; Donadelli,
Alessandro
PATENT ASSIGNEE(S): Pfizer Products Inc., USA
SOURCE: PCT Int. Appl., 47 pp.
CODEN: PIXXD2
DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2002046199	A1	20020613	WO 2001-IB2181	20011119
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM				
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
AU 2002023929	A5	20020618	AU 2002-23929	20011119
US 2002099205	A1	20020725	US 2001-6579	20011204
PRIORITY APPLN. INFO.:			US 2000-251018P	20001204
			WO 2001-IB2181	20011119

OTHER SOURCE(S): CASREACT 137:33162
GI



AB A process for the prepn. of I (R1 = p-nitrobenzyl, allyl; X = halo)

Searcher : Shears 308-4994

via an intramol. Wittig reaction of II (R1 = p-nitrobenzyl, allyl; R2 = C1-6-alkyl, C6-10-aryl, C6-10-aryl-C1-6-alkyl, dithianyl) to prep. 3-cyclic-ether substituted derivs. of cephalosporins is described. Thus, III was treated with p-nitrobenzyl glyoxylate monohydrate followed by redn. of the intermediate with NaBH4. The resulting hydroxy compd. was treated with p-toluenesulfonic acid followed by addn. of (S)-1-(tetrahydro-2-furanyl)ethanone, addn. of thionyl chloride, and finally trimethylphosphine to give the desired intermediate II (R1 = p-nitrobenzyl, R2 = PhCH2). Cyclization of II via an intramol. Wittig reaction was accomplished by refluxing for 16 h in THF. Addn. of phosphorus pentachloride and .alpha.-picoline in dichloromethane gave the free amine of I (R1 = p-nitrobenzyl).

IC ICM C07D501-08
 ICS C07D501-18; C07D501-20; C07D405-12; C07F009-568; C07D205-095;
 C07D513-04; C07D513-04; C07D277-00; C07D205-00

CC 26-5 (Biomolecules and Their Synthetic Analogs)

ST cephalosporin lactam antibiotic cyclic ether substituted prepn;
 Wittig reaction intramol cyclic ether cephalosporin prepn

IT Wittig reaction
 (intramol.; process for the prepn. of p-nitrobenzyl or allyl esters of 3-cyclic-ether substituted cephalosporins from trimethylphosphinic compds. via an intramol. Wittig reaction)

IT Lactams
 RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP (Preparation)
 (.beta.-; process for the prepn. of p-nitrobenzyl or allyl esters of 3-cyclic-ether substituted cephalosporins from trimethylphosphinic compds. via an intramol. Wittig reaction)

IT Antibiotics
 (.beta.-lactam; process for the prepn. of p-nitrobenzyl or allyl esters of 3-cyclic-ether substituted cephalosporins from trimethylphosphinic compds. via an intramol. Wittig reaction)

IT 676-96-0
 (prepn. of)

IT 436100-73-1P 436100-74-2P 436100-75-3P 436100-76-4P
 436100-77-5P 436100-78-6P 436800-38-3P 436800-39-4P
 436800-40-7P
 RL: IMF (Industrial manufacture); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
 (process for the prepn. of p-nitrobenzyl or allyl esters of 3-cyclic-ether substituted cephalosporins from trimethylphosphinic compds. via an intramol. Wittig reaction)

IT 436100-68-4P 436800-42-9P
 RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP (Preparation)
 (process for the prepn. of p-nitrobenzyl or allyl esters of 3-cyclic-ether substituted cephalosporins from trimethylphosphinic compds. via an intramol. Wittig reaction)

IT 64-17-5, Ethanol, uses 67-56-1, Methanol, uses 67-64-1, Acetone, uses 68-12-2, DMF, uses 71-23-8, Propanol, uses 75-09-2, Methylene chloride, uses
 RL: NUU (Other use, unclassified); USES (Uses)
 (process for the prepn. of p-nitrobenzyl or allyl esters of 3-cyclic-ether substituted cephalosporins from trimethylphosphinic compds. via an intramol. Wittig reaction)

IT 79-37-8, Oxalyl chloride 594-09-2, Trimethylphosphine 619-73-8,
 4-Nitrobenzylalcohol 34103-69-0 64370-42-9, Allyl glyoxylate 131328-27-3 141194-61-8 192049-49-3 436800-46-3 436801-05-7

10/006579

436801-06-8 436801-07-9 436801-08-0
RL: RCT (Reactant); RACT (Reactant or reagent)
(process for the prepn. of p-nitrobenzyl or allyl esters of
3-cyclic-ether substituted cephalosporins from
trimethylphosphinic compds. via an intramol. Wittig reaction)
IT 81779-73-9P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation);
RACT (Reactant or reagent)
(process for the prepn. of p-nitrobenzyl or allyl esters of
3-cyclic-ether substituted cephalosporins from
trimethylphosphinic compds. via an intramol. Wittig reaction)
IT 108-48-5, 2,6-Lutidine 109-02-4, N-Methylmorpholine 110-86-1,
Pyridine, reactions 288-32-4, Imidazole, reactions 507-16-4,
Thionyl bromide 7719-09-7, Thionyl chloride 7719-12-2,
Phosphorus trichloride 7789-60-8, Phosphorus tribromide
RL: RGT (Reagent); RACT (Reactant or reagent)
(process for the prepn. of p-nitrobenzyl or allyl esters of
3-cyclic-ether substituted cephalosporins from
trimethylphosphinic compds. via an intramol. Wittig reaction)
REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR
THIS RECORD. ALL CITATIONS AVAILABLE IN
THE RE FORMAT

L14 ANSWER 2 OF 7 MARPAT COPYRIGHT 2002 ACS

(ALL HITs ARE ITERATION INCOMPLETEs)

ACCESSION NUMBER: 136:257270 MARPAT
TITLE: Methods of decreasing or preventing pain using
spicamycin derivatives
INVENTOR(S): Borsook, David
PATENT ASSIGNEE(S): The General Hospital Corporation, USA
SOURCE: PCT Int. Appl., 26 pp.
CODEN: PIXXD2
DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2002024146	A2	20020328	WO 2001-US29371	20010920
W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM				
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				

PRIORITY APPLN. INFO.: US 2000-234382P 20000920

AB Methods of providing pain relief by administering a water-sol.
deriv. of spicamycin. Methods of using pain mediation agents are
also provided.
IC ICM A61K
CC 1-11 (Pharmacology)
ST pain spicamycin analgesia
IT Body, anatomical

10/006579

(back, pain; spicamycin derivs. for prevention and treatment of various pains)
IT Nerve, disease
(diabetic neuropathy; spicamycin derivs. for prevention and treatment of various pains)
IT Drug delivery systems
(implants; spicamycin derivs. for prevention and treatment of various pains)
IT Herpesviridae
(infection, neuropathy; spicamycin derivs. for prevention and treatment of various pains)
IT Drug delivery systems
(injections, i.v.; spicamycin derivs. for prevention and treatment of various pains)
IT Nerve, disease
(neuralgia; spicamycin derivs. for prevention and treatment of various pains)
IT Pancreas, disease
(neuropathy; spicamycin derivs. for prevention and treatment of various pains)
IT Pain
(opioid-resistant; spicamycin derivs. for prevention and treatment of various pains)
IT Viscera
(pain; spicamycin derivs. for prevention and treatment of various pains)
IT Drug delivery systems
(slow-release; spicamycin derivs. for prevention and treatment of various pains)
IT Analgesics
Human
(spicamycin derivs. for prevention and treatment of various pains)
IT 87099-85-2, Spicamycin
RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(spicamycin derivs. for prevention and treatment of various pains)

L14 ANSWER 3 OF 7 MARPAT COPYRIGHT 2002 ACS
(ALL HITs ARE ITERATION INCOMPLETEs)

ACCESSION NUMBER: 135:318706 MARPAT
TITLE: Preparation of halogenated 2-amino-5,6-heptenoic acid derivatives useful as nitric oxide synthase inhibitors
INVENTOR(S): Grapperhaus, Margaret L.; Sikorski, James A.; Awasthi, Alok K.; Wang, Lijuan J.; Pitzele, Barnett S.; Hansen, Donald W., Jr.; Manning, Pamela T.
PATENT ASSIGNEE(S): Pharmacia Corporation, USA
SOURCE: PCT Int. Appl., 133 pp.
CODEN: PIXXD2
DOCUMENT TYPE: Patent
LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
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Searcher : Shears 308-4994

WO 2001078719	A1	20011025	WO 2001-US12258	20010413
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM	RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG			
US 2002049202	A1	20020425	US 2001-835191	20010413
PRIORITY APPLN. INFO.:			US 2000-197032P	20000413

AB Halogenated 2-amino-5,6-heptenoic acid derivs.
R7N:CMeNHCH2CR1:CR2CH2CH2CH(NH2)C(O)J [R1, R2 = H, halo, alkyl,
haloalkyl (at least one of R1 or R2 contains halogen); R7 = H, OH; J
= OH, alkoxy, NR3R4, where R3 = H, alkyl, alkenyl, alkynyl and R4 =
H, (un)substituted heterocyclyl] were prepd. for use as nitric oxide
synthase (NOS) inhibitors. Thus, (2S,5E)-2-amino-6-fluoro-7-[(1-
iminoethyl)amino]-5-heptenoic acid dihydrochloride was prepd. by a
multistep procedure starting from L-glutamic acid and showed IC50
values 0.36, 68, 3.6, and 0.1 .mu.M in hiNOS, hecNOS, hncNOS, and
human cartilage assays, resp.

IC ICM A61K031-221
ICS A61K031-195; A61K031-41; C07C259-14; C07C229-30; C07D271-06;
C07D257-04

CC 34-2 (Amino Acids, Peptides, and Proteins)
Section cross-reference(s): 1, 7

ST acetimidoylaminoheptenoic acid aminohalo prepn inhibitor nitric
oxide synthase; haloaminoheptenoic acid prepn inhibitor nitric oxide
synthase; aminoheptenoic acid halo prepn inhibitor nitric oxide
synthase; heptenoic acid haloamino prepn inhibitor nitric oxide
synthase

IT Alcoholism
Anti-inflammatory agents
Antiarthritis
Antirheumatic agents
Antitumor agents

```
(prepn. of halogenated aminoheptenoic acid derivs. useful as  

nitric oxide synthase inhibitors)
```

IT 54-11-5, Nicotine
RL: BAC (Biological activity or effector, except adverse); BSU
(Biological study, unclassified); BIOL (Biological study)

```
(prepn. of halogenated aminoheptenoic acid derivs. useful as  

nitric oxide synthase inhibitors)
```

IT 367967-68-8P 367967-69-9P 367967-70-2P 367967-71-3P
367967-72-4P 367967-73-5P 367967-74-6P 367967-75-7P
367967-76-8P 367967-77-9P 367967-78-0P 367967-79-1P
367967-80-4P 367967-81-5P 367967-82-6P 367967-83-7P
367967-84-8P 367967-85-9P 367967-86-0P 367967-87-1P
367967-88-2P
RL: BAC (Biological activity or effector, except adverse); BSU
(Biological study, unclassified); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
(Uses)

```
(prepn. of halogenated aminoheptenoic acid derivs. useful as
```

nitric oxide synthase inhibitors)

IT 125978-95-2, Nitric oxide synthase
 RL: BPR (Biological process); BSU (Biological study, unclassified);
 BIOL (Biological study); PROC (Process)
 (prepn. of halogenated aminoheptenoic acid derivs. useful as
 nitric oxide synthase inhibitors)

IT 367968-17-0P
 RL: BYP (Byproduct); PREP (Preparation)
 (prepn. of halogenated aminoheptenoic acid derivs. useful as
 nitric oxide synthase inhibitors)

IT 56-86-0, L-Glutamic acid, reactions 77-76-9, 2,2-Dimethoxy propane
 401-56-9, Ethyl chlorofluoroacetate 696-63-9, p-
 Methoxybenzenethiol 1000-84-6, Ethyl acetimidate 1074-82-4,
 Potassium phthalimide 1499-55-4, L-Glutamic acid 5-methyl ester
 2356-16-3 4418-61-5, 5-Aminotetrazole 52386-40-0
 RL: RCT (Reactant); RACT (Reactant or reagent)
 (prepn. of halogenated aminoheptenoic acid derivs. useful as
 nitric oxide synthase inhibitors)

IT 45214-91-3P 59279-60-6P 126587-35-7P 129599-92-4P
 129600-92-6P 136904-77-3P 144090-56-2P 192314-71-9P
 206128-03-2P 367967-89-3P 367967-90-6P 367967-91-7P
 367967-92-8P 367967-93-9P 367967-94-0P 367967-95-1P
 367967-96-2P 367967-97-3P 367967-98-4P 367967-99-5P
 367968-00-1P 367968-01-2P 367968-02-3P 367968-03-4P
 367968-04-5P 367968-05-6P 367968-06-7P 367968-07-8P
 367968-08-9P 367968-09-0P 367968-10-3P 367968-11-4P
 367968-12-5P 367968-13-6P 367968-14-7P
 RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation);
 RACT (Reactant or reagent)
 (prepn. of halogenated aminoheptenoic acid derivs. useful as
 nitric oxide synthase inhibitors)

REFERENCE COUNT: 8 THERE ARE 8 CITED REFERENCES AVAILABLE FOR
 THIS RECORD. ALL CITATIONS AVAILABLE IN
 THE RE FORMAT

L14 ANSWER 4 OF 7 MARPAT COPYRIGHT 2002 ACS
 (ALL HITs ARE ITERATION INCOMPLETEs)

ACCESSION NUMBER: 134:366693 MARPAT

TITLE: Preparation of bis(aminoalkyl- or
 amidinophenoxy)arylene- and heteroatom-
 interrupted alkanes and analogs as tryptase
 inhibitors

INVENTOR(S): Anderskewitz, Ralf; Braun, Christine; Hamm,
 Rainer; Disse, Bernd; Jennewein, Hans Michael;
 Speck, Georg

PATENT ASSIGNEE(S): Boehringer Ingelheim Pharma K.-G., Germany

SOURCE: Ger. Offen., 36 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent

LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 19955476	A1	20010523	DE 1999-19955476	19991118
WO 2001036374	A2	20010525	WO 2000-EP11216	20001114
WO 2001036374	A3	20020411		

W: AE, AU, BG, BR, CA, CN, CZ, EE, HU, ID, IL, IN, JP, KR, LT,
 LV, MX, NO, NZ, PL, RO, SG, SI, SK, UA, US, UZ, VN, YU, ZA,
 AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
 RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC,
 NL, PT, SE, TR

PRIORITY APPLN. INFO.: DE 1999-19955476 19991118

AB B1Z1X1Z2X2ZX3Z3X4Z4B2 [I; B1,B2 = C(:NR1)NHR1', CH2NH2, CH2CH2NH2,
 ureido; R1,R1' = OH, COR2, CO2R2; R2 = H, alkyl, aryl(alkyl); X1-X4
 = bond, CH2, CH2CH2, CH2O, CH2NH, etc.; Z = (heteroatom-
 interrupted)alkylene, G1(CH2)rG2 [X2 or X3 = (CH2)1-2], E1(CH2)rE2,
 etc.; E1,E2 = azacycloalkylene; G1,G1 = bond or cycloalkylene; Z1-Z4
 = (un)substituted (hetero)arylene; r = 0-6] were prepd. Thus,
 3-(C1H2C)C6H4CH2OC6H4(CH2CH2NHBOc)-4 was condensed with
 (CH2CMe2NH2)2 to give, after deprotection, the N,N'-bisbenzylated
 hexandiamine.4HCl. Data for biol. activity of I were given.

IC ICM C07C217-58
 ICS C07C217-60; C07C213-02; C07D211-26; C07D295-12; C07C257-18;
 C07C259-10; C07C271-62; C12N009-99; A61K031-155

CC 25-19 (Benzene, Its Derivatives, and Condensed Benzenoid Compounds)
 Section cross-reference(s): 1

ST azaarylenealkane bisaminophenoxy bisamidinophenoxy prepn tryptase
 inhibitor; antiinflammatory azaarylenealkane bisaminophenoxy
 bisamidinophenoxy prepn; antiallergic azaarylenealkane
 bisaminophenoxy bisamidinophenoxy prepn

IT Allergy inhibitors
 Anti-inflammatory agents
 (prepn. of bis(aminoalkyl- or amidinophenoxy)arylene- and
 heteroatom-interrupted alkanes and analogs as tryptase
 inhibitors)

IT 97501-93-4, Tryptase
 RL: BPR (Biological process); BSU (Biological study, unclassified);
 BIOL (Biological study); PROC (Process)
 (mediated disorders; treatment; prepn. of bis(aminoalkyl- or
 amidinophenoxy)arylene- and heteroatom-interrupted alkanes and
 analogs as tryptase inhibitors)

IT 340284-41-5P 340284-43-7P 340284-44-8P 340284-45-9P
 340284-46-0P 340284-48-2P 340284-49-3P 340284-50-6P
 340284-51-7P 340284-52-8P 340284-53-9P 340284-54-0P
 340284-55-1P 340284-56-2P 340284-57-3P 340284-58-4P
 340284-59-5P 340284-60-8P 340284-61-9P 340284-62-0P
 340284-63-1P 340284-64-2P 340284-65-3P 340284-66-4P
 340284-67-5P 340284-68-6P 340284-69-7P 340284-70-0P
 340284-71-1P 340284-72-2P 340284-73-3P 340284-74-4P
 340284-75-5P 340284-76-6P 340284-77-7P 340284-78-8P
 340284-79-9P 340284-80-2P 340284-81-3P 340284-82-4P
 340284-83-5P 340284-84-6P 340284-85-7P 340284-86-8P
 340284-90-4P 340284-91-5P 340284-92-6P 340284-93-7P
 340284-94-8P 340284-95-9P 340284-96-0P 340284-97-1P
 340284-98-2P 340284-99-3P 340285-00-9P 340285-01-0P
 340285-02-1P 340285-03-2P 340285-04-3P 340285-05-4P
 340285-06-5P 340285-07-6P 340285-09-8P 340285-10-1P
 340285-11-2P 340285-12-3P 340285-13-4P 340285-14-5P
 340285-15-6P 340285-16-7P

RL: BAC (Biological activity or effector, except adverse); BSU
 (Biological study, unclassified); SPN (Synthetic preparation); THU
 (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
 (Uses)
 (prepn. of bis(aminoalkyl- or amidinophenoxy)arylene- and

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heteroatom-interrupted alkanes and analogs as tryptase
inhibitors)
IT 23578-35-0, 2,5-Diamino-2,5-dimethylhexane 255915-70-9
340284-87-9 340284-88-0 340284-89-1
RL: RCT (Reactant); RACT (Reactant or reagent)
(prep. of bis(aminoalkyl- or amidinophenoxy)arylene- and
heteroatom-interrupted alkanes and analogs as tryptase
inhibitors)

L14 ANSWER 5 OF 7 MARPAT COPYRIGHT 2002 ACS
(ALL HITs ARE ITERATION INCOMPLETEs)
ACCESSION NUMBER: 131:5098 MARPAT
TITLE: Acylation of aromatic compounds
INVENTOR(S): Baudry, Barbier Denise; Dormond, Alain; Richard,
Stephanie; Desmurs, Jean Roger
PATENT ASSIGNEE(S): Rhodia Chimie, Fr.
SOURCE: Fr. Demande, 38 pp.
CODEN: FRXXBL
DOCUMENT TYPE: Patent
LANGUAGE: French
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
FR 2768729	A1	19990326	FR 1997-11701	19970919

OTHER SOURCE(S): CASREACT 131:5098
AB The title process takes place in the presence of, e.g., a rare earth halide. Thus, benzoylation of anisole gave 90.3% 4-(MeO)C₆H₄COPh in the presence of NdCl₃.dioxane.
IC ICM C07C049-76
ICS C07C045-45; B01J027-125; B01J031-22
ICI B01J031-22, B01J103-26
CC 25-16 (Benzene, Its Derivatives, and Condensed Benzenoid Compounds)
ST acylation arom rare earth catalyst; benzophenone prepn
IT Ketones, preparation
RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP (Preparation)
(arom.; acylation of arom. compds.)
IT Acylation catalysts
Acylation catalysts
(benzoylation catalysts; acylation of arom. compds.)
IT Benzoylation
Benzoylation
(catalysts; acylation of arom. compds.)
IT 611-94-9P, p-Methoxybenzophenone 5672-94-6P, 1-Acetyl-2-methoxynaphthalene 5703-21-9P, 4-Acetylveratrole
RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP (Preparation)
(acylation of arom. compds.)
IT 91-16-7, Veratrole 93-04-9, 2-Methoxynaphthalene 98-88-4, Benzoyl chloride 100-66-3, Anisole, reactions
RL: RCT (Reactant); RACT (Reactant or reagent)
(acylation of arom. compds.)
IT 10024-93-8, Neodymium trichloride 10361-92-9, Yttrium trichloride
RL: CAT (Catalyst use); USES (Uses)
(catalyst for acylation of arom. compds.)

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L14 ANSWER 6 OF 7 MARPAT COPYRIGHT 2002 ACS
(ALL HITs ARE ITERATION INCOMPLETEs)

ACCESSION NUMBER: 131:5097 MARPAT
TITLE: Acylation of aromatic compounds
INVENTOR(S): Baudry, Barbier Denise; Dormond, Alain; Richard,
Stephanie; Bouazza, Aicha; Desmurs, Jean Roger
PATENT ASSIGNEE(S): Rhodia Chimie, Fr.
SOURCE: Fr. Demande, 28 pp.
CODEN: FRXXBL
DOCUMENT TYPE: Patent
LANGUAGE: French
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
FR 2768728	A1	19990326	FR 1997-11700	19970919
FR 2768728	B1	19991203		

OTHER SOURCE(S): CASREACT 131:5097
AB The title process takes place in the presence of a U or uranyl halide. Thus, benzoylation of anisole gave 93% 4-(MeO)C₆H₄COPh after 1h reflux in the presence of a catalyst prep'd. from U308 and HCl.
IC ICM C07C049-76
 ICS C07C049-786; C07C049-84; B01J027-08
ICI B01J027-08, B01J103-28
CC 25-16 (Benzene, Its Derivatives, and Condensed Benzenoid Compounds)
ST acylation arom uranium catalyst; benzophenone prep'n
IT Ketones, preparation
 RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP (Preparation)
 (arom.; acylation of arom. compds.)
IT Acylation catalysts
 Acylation catalysts
 (benzoylation catalysts; acylation of arom. compds.)
IT Benzoylation
 Benzoylation
 (catalysts; acylation of arom. compds.)
IT 134-84-9P 611-94-9P 954-16-5P 4044-60-4P 4885-14-7P
 6317-73-3P 26086-67-9P 40777-50-2P 225780-54-1P 225780-55-2P
 RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP (Preparation)
 (acylation of arom. compds.)
IT 95-93-2, Durene 98-88-4, Benzoyl chloride 100-20-9,
 1,4-Benzenedicarbonyl dichloride 100-66-3, Anisole, reactions
 101-84-8, Diphenyl oxide 106-42-3, p-Xylene, reactions 108-67-8,
 Mesitylene, reactions 108-88-3, Toluene, reactions
 RL: RCT (Reactant); RACT (Reactant or reagent)
 (acylation of arom. compds.)

L14 ANSWER 7 OF 7 MARPAT COPYRIGHT 2002 ACS
(ALL HITs ARE ITERATION INCOMPLETEs)

ACCESSION NUMBER: 130:196650 MARPAT
TITLE: 2-Benzoylcyclohexane-1,3-diones as herbicides
INVENTOR(S): Engel, Stefan; Rheinheimer, Joachim; Baumann,
Ernst; Von Deyn, Wolfgang; Hill, Regina Luise;
Mayer, Guido; Misslitz, Ulf; Wagner, Oliver;
Witschel, Matthias; Otten, Martina; Walter,

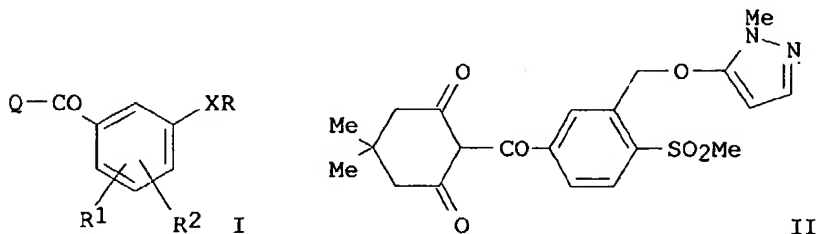
Searcher : Shears 308-4994

10/006579

PATENT ASSIGNEE(S): Helmut; Westphalen, Karl-otto; et al
SOURCE: BASF Aktiengesellschaft, Germany
PCT Int. Appl., 81 pp.
CODEN: PIIXXD2
DOCUMENT TYPE: Patent
LANGUAGE: German
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 9910327	A1	19990304	WO 1998-EP4634	19980805
W: AL, AU, BG, BR, BY, CA, LT, LV, MK, MX, NO, NZ, VN, AM, AZ, BY, KG, KZ,	CN, CZ, GE, HU, ID, IL, JP, KR, KZ, PL, RO, RU, SG, SI, SK, TR, UA, US,			
RW: AT, BE, CH, CY, DE, DK, NL, PT, SE	MD, RU, TJ, TM	ES, FI, FR, GB, GR, IE, IT, LU, MC,		
AU 9890684	A1	19990316	AU 1998-90684	19980805
EP 1001938	A1	20000524	EP 1998-942611	19980805
R: CH, DE, FR, GB, LI				
JP 2001514171	T2	20010911	JP 2000-507656	19980805
ZA 9807057	A	20000207	ZA 1998-7057	19980806
US 6432881	B1	20020813	US 2000-485231	20000207
PRIORITY APPLN. INFO.:			DE 1997-19734164	19970807
			WO 1998-EP4634	19980805

GI



AB The 2-benzoylcyclohexane-1,3-diones I [Q = (un)substituted 1,3-dioxo-2-cyclohexyl; X = nalkylene, oxaalkylene, thiaalkylene; R = heterocyclic; R₁, R₂ = H, NO₂, halogen, CN, SCN, (un)substituted alkyl, OH, SH, SO₃H, SO₂NH₂, NHSO₂H, acylamino] were prep'd. for use as herbicides (no data). Thus, Me 2-chloro-3-methyl-4-methylsulfonylbenzoate was treated with 1-methyl-5-pyrazolol, hydrolyzed to the acid and treated with dimedone to give the benzoylpyrazole II.

IC ICM C07D231-12
ICS C07D231-14; A01N043-56

CC 28-8 (Heterocyclic Compounds (More Than One Hetero Atom))
Section cross-reference(s): 5

ST benzoylcyclohexanedione prepn herbicide;
pyrazolylbenzoylcyclohexanedione prepn herbicide;
pyridylbenzoylcyclohexanedione prepn herbicide

IT Herbicides
(prepn. of benzoylcyclohexanediones as herbicides)

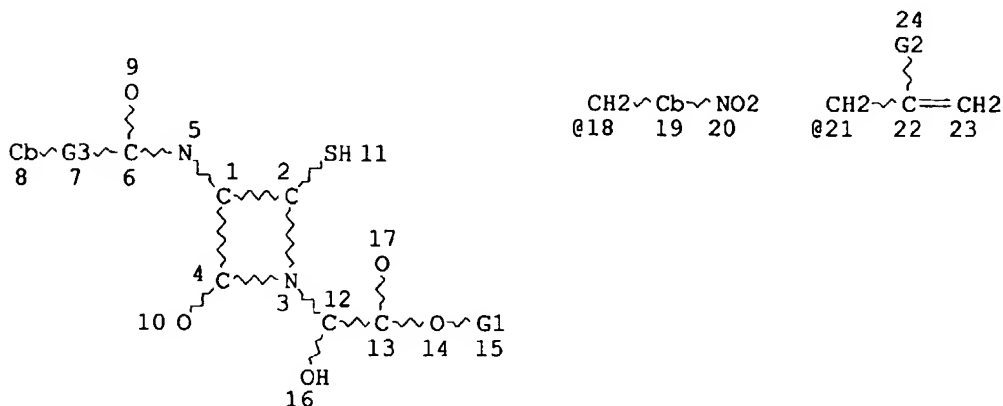
IT 220798-99-2P 220799-06-4P 220799-10-0P 220799-15-5P

Searcher : Shears 308-4994

10/006579

220799-18-8P 220799-24-6P 220799-29-1P 220799-33-7P
RL: AGR (Agricultural use); SPN (Synthetic preparation); BIOL
(Biological study); PREP (Preparation); USES (Uses)
(prepn. of benzoylcyclohexanediones as herbicides)
IT 126-81-8, Dimedone 33641-15-5, 5-Hydroxy-1-methylpyrazole
120100-04-1
RL: RCT (Reactant); RACT (Reactant or reagent)
(prepn. of benzoylcyclohexanediones as herbicides)
IT 120100-44-9P 220798-89-0P 220798-93-6P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation);
RACT (Reactant or reagent)
(prepn. of benzoylcyclohexanediones as herbicides)
REFERENCE COUNT: 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR
THIS RECORD. ALL CITATIONS AVAILABLE IN
THE RE FORMAT

L7 FOLIO 'MARPATPREV' ENTERED AT 15:49:55 ON 04 OCT 2002
STR



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VAR G1=18/21
VAR G2=H/CH3
REP G3=(1-6) C
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
GGCAT IS UNS AT 8
DEFAULT ECLEVEL IS LIMITED
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GRAPH ATTRIBUTES:
RSPEC I
NUMBER OF NODES IS 24

STEREO ATTRIBUTES: NONE

ATTRIBUTES SPECIFIED AT SEARCH-TIME:
MLEVEL IS CLASS ON RING NODES AND RING GROUPS
MLEVEL IS CLASS ON CHAIN NODES AND CHAIN GROUPS.
ECLEVEL IS LIM ON ALL NODES
ALL RING(S) ARE ISOLATED

0-SEA FILE=MARPATPREV SSS FUL L7 (MODIFIED ATTRIBUTES)

100.0% PROCESSED 127 ITERATIONS 0 ANSWERS

Searcher : Shears 308-4994

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SEARCH TIME: 00.00.06

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FILE 'HOME' ENTERED AT 15:50:21 ON 04 OCT 2002

Searcher : Shears 308-4994